IN THE CLAIMS

Please cancel claims 1-2, 6-7 and 10-15. Please amend claims 3, 5, and

8 and add new claims 16-21 to provide as follows.

(Currently Amended) A viable cellular population produced by a method

comprising infecting the cells with a retrovirus in the presence of an effective

immobilized amount of material including a ligand which binds to the cells and a

ligand which binds to the retrovirus, so as to co-localize the retrovirus and the cells

and increase the transduction efficiency of the cells, said infecting being conducted

in a medium essentially free from hexadimethrine bromide.

(Original) The viable cellular population of claim 3 which comprises 4.

hematopoietic stem cells.

5. (Currently Amended) A method for cellular grafting, comprising:

grafting a mammal with a viable cellular population produced by a method

comprising infecting the cells with a retrovirus in the presence of an effective

immobilized amount of material including a ligand which binds to the cells and a

ligand which binds to the retrovirus, so as to co-localize the retrovirus and the cells

and increase the transduction efficiency of the cells, said infecting being conducted

in a medium essentially free from hexadimethrine bromide.

8. (Currently Amended) A method for cellular grafting, comprising:

grafting a mammal with a cellular composition, comprising a substantially retroviral-

tranduced population of viable cells, said composition being essentially free from

both retroviral producer cells and hexadimethrine bromide.

9. (Original) The method of claim 8 wherein the cellular population comprises

hematopoietic stem cells.

16. (New) The method of claim 3, wherein the medium is essentially free from any polycationic agent which increases the efficiency of transduction of the viable mammalian cells by the retrovirus in co-culture, but which agent reduces

the efficiency of transduction of the cells by the retrovirus in the presence of said

material.

17. (New) The method of claim 16, wherein said material comprises

substantially pure fibronectin, substantially pure fibronectin fragments, or mixture

thereof.

18. (New) The method of claim 5, wherein the medium is essentially free

from any polycationic agent which increases the efficiency of transduction of the

viable mammalian cells by the retrovirus in co-culture, but which agent reduces

the efficiency of transduction of the cells by the retrovirus in the presence of said

material.

19. (New) The method of claim 18, wherein said material comprises

substantially pure fibronectin, substantially pure fibronectin fragments, or mixture

thereof.

20. (New) The method of claim 8, wherein said composition is essentially

free from any polycationic agent that is effective to increase the efficiency of

transduction of the viable cells by the retrovirus in co-culture.

21. (New) The method of claim 20, wherein said viable cells have been

transduced in the presence of substantially pure fibronectin, substantially pure

fibronectin fragments, or mixture thereof, so as to increase the efficiency of

transduction by the retrovirus.

Preliminary Amendment Inventor: Williams, David A.